Enol Esters. III. Preparation of Disopropenyl Esters of Dicarboxylic Acids

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In previous publications we have reported the preparation of isopropenyl esters⁴ of monocarboxylic, long-chain fatty acids and have shown their usefulness as acylating agents.⁵ Acylation occurs even in many unfavorable cases because the evolution of gaseous acetone tends to drive the reaction essentially to completion; for example, N-alkylamides and imidamides

For the previous paper in this series, see E. S. Rothman, S. Serota, and
Swern, J. Org. Chem., 29, 646 (1964).

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(4) E. S. Rothman, S. Serota, T. Perlstein, and D. Swern, J. Org. Chem. 27, 3123 (1962).

(5) E. S. Rothman, S. Serota, and D. Swern, ibid., 29, 646 (1964).

such as barbituric acid acylate easily. Not only are clean reaction products obtained in a simple operation, but there is no problem of removing pyridine or emulsified metal soaps as is the case when conventional reagents such as stearoyl chloride or steric anhydride are used.

It occurred to us that by extending this acylation principle to bi- or polyfunctional materials one would have at hand the basis for several polymer-forming reactions.6 To follow this line, we decided to prepare the difunctional acylating agents, diisopropenyl azelate, diisopropenyl sebacate, and related substances. This goal was not so easy to accomplish as anticipated. At first we were baffled by the apparently great difference in behavior between the monocarboxylic and dicarboxylic acids in isopropenyl ester formation, but gradually we were able to proceed from small yields to useful yields and present a summary of the properties of the compounds prepared in Table I. It is the purpose of the present paper to explain the reasons for the peculiar behavior and to indicate how the deleterious actions can be circumvented by improved techniques.

(6) E. S. Rothman, Tetrahedron Letters, in press.

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	TABLE I			
•	Bp (mm)			
	or			Yield,
Diisopropenyl ester	mp, °C	n 30 D	d^{30}_{4}	%
Sebacate	140 (0.13)	1.4488	0.9698	80-85
Azelate	103 (0.03)	1.4499	0.9746	75
Suberate	94 (0.025)	1.4470	0.9775	70
Dodecanedioate		1.4505	0.9571	40
Tridecanedioate		1.4509	0.9464	40
Tetradecanedioate	31.5-33			40
Adipate	88 (0.11)	1.4453	0.9826	70
Isopropenyl hydro- gen sebacate	40-41		- · ·	• •

There are two main routes that we have studied in disopropenyl dioate formation. The first involves the direct addition of 1 mole of dioic acid to 2 moles of propyne. The second route involves the acidester interchange reaction, utilizing the commercially available isopropenyl acetate, in excess, as the isopropenylating agent. In special cases one procedure may give better yields than the other; for example, the propyne procedure may be preferable for conversions of α-halogenated acids, but in general the problem to be solved is not that of forming the product in useful yield, but rather that of isolation of the product without destroying it during the isolation procedure. That is to say, the formation of diisopropenyl ester itself offers no unusual problems, but so many side reactions can carry the already formed product to undesirable material that the yield may be dropped to very small values if these side reactions are allowed to take place.

The major sort of side reaction involves the addition of the intermediate half-ester, e.g., isopropenyl hydrogen sebacate, at the carboxylic acid end to the isopropenyl ester group of another molecule. The resulting unstable geminal diester rapidly collapses, expelling acetone and forming a "dimeric" molecule with an anhydride connecting linkage. The process may be repeated several times and followed by observing the changing infrared carbonyl absorption spectrum. As time progresses, the ester bands increase in intensity to a maximum and then disappear as anhydride bands increase in intensity concurrently. During the work-up, because of the difficulty in removing the catalyst completely, the point in time at which ester function is at a maximum is reached and passed until, sometimes, only anhydride is isolable. For good diester yields it is necessary to freeze all equilibria prior to work-up. Use of salts in heterogeneous phase, such as potassium acetate or potassium bicarbonate, or liquid bases, such as tertiary amines, is helpful in quenching but is not completely reliable.

Absorptive removal through a short clean-up column of Florisil not only removes catalyst and gives clarification of suspended matter including a purple coloration that appears to be colloidal metallic mercury owing to reduction of mercury salts in propyne runs, but also removes a small proportion of contaminant unreacted acid and anhydride. Such a preliminary "chromatography" seems to be innocuous, but standard chromatography procedures using a high ratio of adsorbent to compound are risky. Long-chain iso-

propenyl esters such as isopropenyl stearate (C₁₈) may be chromatographed, but even isopropenyl laurate (C₁₂) is largely destroyed. In general, the longer the methylene chain the safer the chromatographic procedures. To rationalize its high lability. one may consider that isopropenyl sebacate (C₁₀) behaves like a pair of pentanoic acid molecules. Sometimes the heat of reaction with the column support is so great that the column feels quite hot to the touch.

After such preliminary passage through a short cleanup column, the recommended procedure is a preliminary distillation in a falling-film molecular distillation apparatus. The time of actual heating during such distillation is kept minimal, so that little pyrolysis or artifact anhydride formation occurs. Curiously enough, the appropriate molecular distillate fractions may now be redistilled all day long through a 24-in. spinning-band column with no perceptible decomposition. Attempts to use such spinning-band apparatus directly on crude reaction products occasionally succeed, but, more typically, results are ruinous.

In passing we should like also to call attention to the fact that treatment of the mixture of dioic acid, mercuric salts, and boron trifluoride etherate in methylene chloride solution with propyne always causes some reduction of mercury salts to free mercury, calling to mind analogous reductions of palladium salts by ethylene.8

Experimental Section

Diisopropenyl Sebacate. Procedure A (Acid-Ester Interchange).—Sebacic acid (202 g, 1 mole), isopropenyl acetate (1.1 l., 11 moles), mercuric acetate (10.0 g, 0.03 mole), and boron trifluoride etherate (4.6 ml) were mixed at 15° and allowed to warm to ambient temperature of 25°. In 2 hr infrared analysis showed total disappearance of the carboxyl carbonyl band. The mixture was treated with 4.1 ml of triethylamine and concentrated in vacuo at 30° to remove most of the excess isopropenyl acetate. The residue was diluted with pentane and filtered through 190 g of Florisil in a short column. The column was rinsed with pentane, and the two percolates were combined. After removing the pentane, the residue was distilled in a fallingfilm molecular still. The fraction collected at about 50 μ and 100-110° was the required diisopropenyl sebacate (155 g). On redistillation through a 24-in. spinning-band column, a high recovery of diisopropenyl sebacate was obtained: bp 134° (0.06-0.07 mm), 140° (0.13 mm); yield 146 g; n^{30} D 1.4488; $d^{30}_{4} = 0.9698$; $\nu_{\text{max}}^{\text{CS2}}$ 1751 (CO) and 1673 (C=C) cm⁻¹. Anal. Calcd for $C_{16}H_{24}O_{4}$: C, 68.05; H, 9.28. Found:

C, 68.04; H, 9.22.

Diisopropenyl Azelate. Procedure B (Addition of Acid to Propyne).—Azelaic acid (100 g, 0.53 mole), methylene chloride (500 ml), and mercuric acetate (18.2 g, 0.057 mole) were treated with 1 ml of boron trifluoride etherate, and propyne gas (freed of anticatalytic matter by passing through Linde Co. Molecular Sieve 4A) was bubbled through the mixture for 3 hr. reaction, once started, was exothermic and was controlled by the rate of gas feed and/or by the cooling owing to the low boiling point of the methylene chloride solvent. Solid sodium bicarbonate (5 g) was added and the mixture was filtered free of solids and metallic mercury. The filtrate was passed through a short Florisil column which was then washed with additional methylene chloride. The residue obtained by evaporation of the percolate was then submitted to molecular distillation and spinning-band column distillation similar to the operations noted in procedure A. The product boiled at 103° (0.03 mm): n^{30} D 1.4499; d^{30}_4 0.9746; $\nu_{\text{max}}^{\text{CS}_2}$ 1755, 1675, 1376, 1200, 1150, and 867 cm⁻¹.

Anal. Calcd for C15H24O5: C, 67.13; H, 9.02. Found: C 67.68; H, 9.25.

⁽⁷⁾ Cf. R. L. Adelman, J. Org. Chem., 14, 1057 (1949), and references cited

⁽⁸⁾ J. Smidt, W. Hafner, R. Jira, R. Sieber, J. Sedlmeier, and A. Sabel Angew. Chem., 74, 93 (1962).